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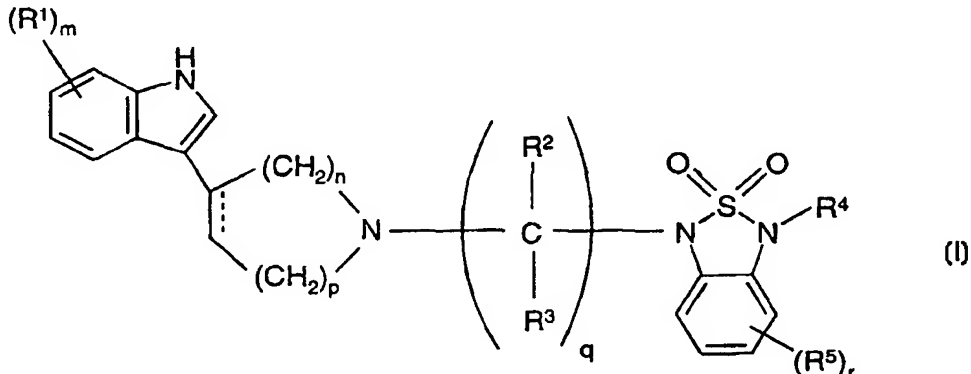


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(54) Title: **1-((INDOLYL AZACYCLOALKYL) ALKYL)-2,1,3-BENZOTHIADIAZOLE 2,2-DIOXIDES EXHIBITING 5-HT2A RECEPTOR ACTIVITY**



(57) Abstract

A pharmaceutical compound of (I) in which m is 0, 1 or 2, n is 1 or 2, p is 1 or 2, q is 1 to 6, r is 0 or 1 to 3, R¹ is halo, C₁₋₄ alkyl, nitrile, trifluoromethyl or C₁₋₄ alkoxy, R² and R³ are each hydrogen or C₁₋₄ alkyl, R⁴ is hydrogen, C₁₋₄ alkyl, optionally substituted phenyl or optionally substituted phenyl-C₁₋₄ alkyl, R⁵ is C₁₋₄ alkyl, C₁₋₄ alkoxy, carboxy, hydroxy, cyano, halo, trifluoromethyl, nitro or amino, and the dotted line represents an optional double bond; and salts and esters thereof.

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1-((INDOLYL AZACYCLOALKYL) ALKYL)-2,1, 3-BENZOTHIADIAZOLE 2,2-DIOXIDES EXHIBITING 5-HT2A RECEPTOR ACTIVITY

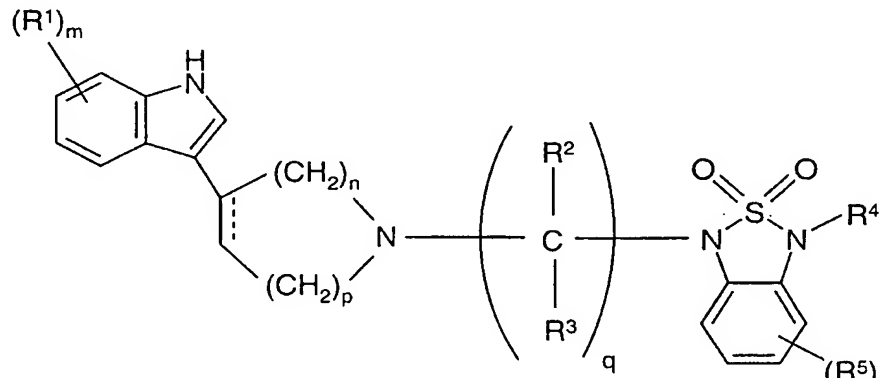
This invention relates to novel compounds with pharmaceutical properties.

5

It is well known that compounds active at serotonin receptors have potential in the treatment of disorders of the central nervous system and, for example, certain halo-substituted indole compounds having serotonin 10 antagonist properties are disclosed in EP-A 0433149 and WO 98/31686.

The compounds of the invention are of the following formula:

15



in which m is 0, 1 or 2, n is 1 or 2, p is 1 or 2,

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q is 1 to 6, r is 0 or 1 to 3,

R¹ is halo, C₁₋₄ alkyl, nitrile, trifluoromethyl or C₁₋₄ 5 alkoxy,

R² and R³ are each hydrogen or C₁₋₄ alkyl,

R⁴ is hydrogen, C₁₋₄ alkyl, optionally substituted 10 phenyl or optionally substituted phenyl-C₁₋₄ alkyl,

R⁵ is C₁₋₄ alkyl, C₁₋₄ alkoxy, carboxy, hydroxy, cyano, 15 halo, trifluoromethyl, nitro or amino, and the dotted line represents an optional double bond;

and salts and esters thereof.

The compounds of the invention and their 20 pharmaceutically acceptable salts and esters are indicated for use in the treatment of disorders of the central nervous system.

A C₁₋₄ alkyl group can be methyl, ethyl or propyl and 25 can be branched or unbranched and includes isopropyl and

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tert. butyl. A C₁₋₄ alkoxy group is one such C₁₋₄ alkyl group attached through oxygen to the ring. An optionally substituted phenyl-C₁₋₄ alkyl group is an optionally substituted phenyl attached through one such 5 C₁₋₄ alkyl group, and is preferably optionally substituted phenyl-(CH₂)_x- where x is 1 or 2, and most preferably optionally substituted benzyl. A halo substituent is preferably fluoro, chloro or bromo.

10 An optionally substituted phenyl group is optionally substituted with one or more, preferably one to three, substitutents selected from, for example C₁₋₄ alkyl, C₁₋₄ alkoxy, carboxy, hydroxy, cyano, halo, trifluoromethyl, nitro and amino.

15

It will be appreciated that when m is 2 the values of R¹ need not be the same, when p is more than one, the recurring unit is not necessarily the same, and when r is 2 or 3 the values of R⁵ need not be the same.

20

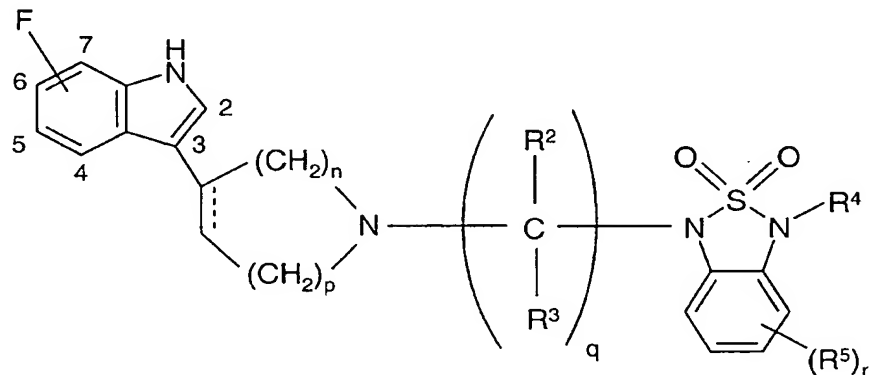
A preferred group of compounds is one of formula (I) above, in which the dotted line represents a double bond, n is 2, m is 1 or 2, and p is 1, R² and R³ are both hydrogen, q is 2, R³ is C₁₋₄ alkyl, and r is 0 or

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1. When R^1 is C_{1-4} alkyl or C_{1-4} alkoxy it is
preferably methyl or methoxy, respectively.

Substituent $(R^1)_m$ preferably represents 6-fluoro,
5 7-fluoro, 6,7-difluoro, or 6-fluoro-7-methyl.

Preferred compounds are those of the following
formula (II):



10

(II)

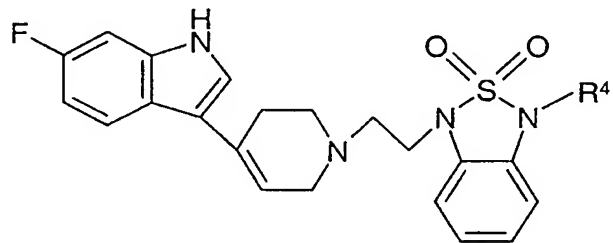
and preferred sub-groups exhibit one or more of the
following features:

15 (i) the fluorine substituent is in the 6- or
7-position, and preferably in the 6-position
(ii) the dotted line represents a double bond
(iii) n is 2 and p is 1

- 5 -

- (iv) R² and R³ are both hydrogen
- (v) q is 2
- (vi) R⁴ is C₁₋₄ alkyl, especially isopropyl
- (vii) r is 0 or 1, and preferably 0 (unsubstituted)
- 5 (viii) R⁵ is C₁₋₄ alkoxy, hydroxy, halo or amino (-NH₂).

A particularly preferred group of compounds is of the formula:



10

(III)

in which R⁴ is C₁₋₄ alkyl and especially isopropyl, or a pharmaceutically acceptable salt thereof.

- 15 As indicated above, it is, of course, possible to prepare salts of the compound of the invention and such salts are included in the invention. Acid addition salts are preferably the pharmaceutically acceptable, non-toxic addition salts with suitable acids, such as
- 20 those with inorganic acids, for example hydrochloric,

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hydrobromic, nitric, sulphuric or phosphoric acids, or with organic acids, such as organic carboxylic acids, for example, glycollic, maleic, hydroxymaleic, fumaric, malic, tartaric, citric, salicyclic, o-acetoxybenzoic, 5 or organic sulphonic, 2-hydroxyethane sulphonic, toluene-p-sulphonic, or naphthalene-2-sulphonic acid.

In addition to the pharmaceutically acceptable salts, other salts are included in the invention. They may 10 serve as intermediates in the purification of compounds or in the preparation of other, for example pharmaceutically acceptable, acid addition salts, or are useful for identification, characterisation or purification.

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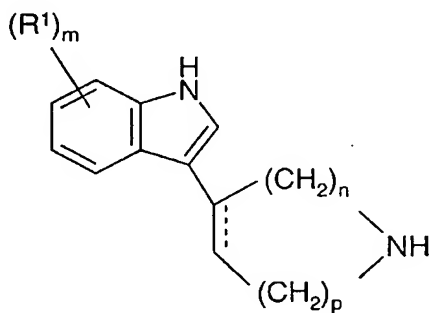
It will be appreciated that when a phenyl substituent is acidic such as, for example, a carboxy group, the opportunity exists for esters. These can be aliphatic or aromatic, being preferably alkyl esters derived from 20 C₁₋₄ alkanols, especially methyl and ethyl esters. An example of an ester substituent is -COOR' where R' is C₁₋₄ alkyl.

Some of the compounds of the invention contain one or 25 more asymmetric carbon atoms which gives rise to isomers. These compounds are normally prepared as

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racemic mixtures and can conveniently be used as such, but individual isomers can be isolated by conventional techniques, if so desired. Such racemic mixtures and individual optical isomers form part of the present 5 invention. It is preferred to use an enantiomerically pure form.

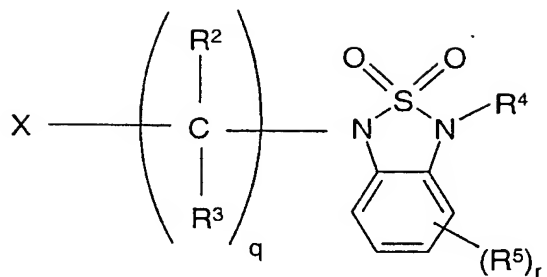
The invention also includes a process for producing a compound of formula (I) above, which comprises reacting 10 a compound of the formula:



(IV)

with a compound of the formula:

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(V)

where the substituents have the values given above, and X is a leaving group such as, for example, a halo atom, 5 or a mesylate or tosylate.

The reaction is preferably carried out in a polar solvent such as, for example, acetonitrile or water, at a temperature of from 50° C. to 150° C., and in the 10 presence of sodium iodide and a base such as, for example, sodium carbonate

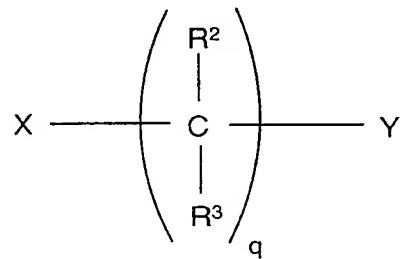
The coupling can also be effected by reacting the compound of formula (IV) with an aldehyde equivalent of 15 the compound of formula (V). Such aldehydes can be prepared from the appropriate terminal alkene by oxidation employing, for example, ozone or osmium tetroxide, followed by reductive amination using, for example, sodium cyanoborohydride, borane in pyridine or 20 sodium triacetoxy borohydride, and the compound of

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formula (IV). This reaction is preferably carried out at a temperature of from -20° C. to 50° C., in a solvent such as, for example, dichloromethane.

5 The intermediate compounds of formula (IV) are known in the art, and compounds of formula (V) can be prepared by preparative routes as follows. For example, compounds of formula (V) can be prepared by reacting the appropriate alkane derivative of formula:

10

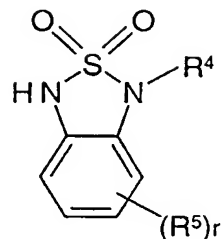


(VI)

where X is a leaving group, and Y is halo, preferably bromo, with a compound of formula:

15

- 10 -



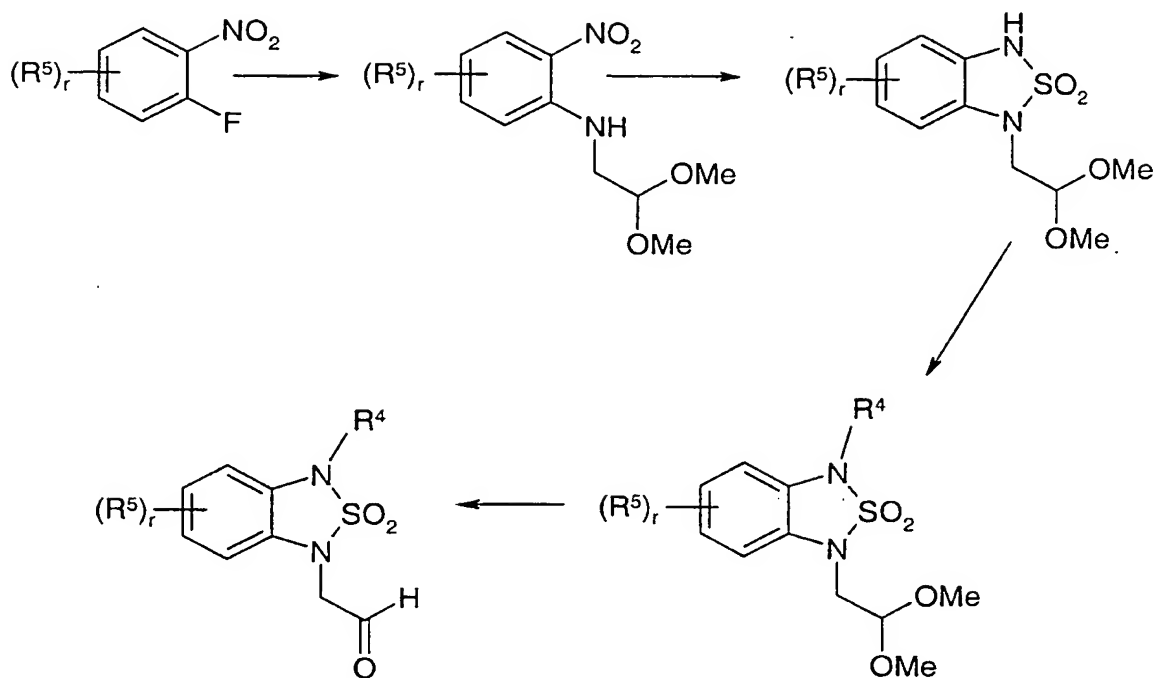
(VII)

Preferred alkane reactants are dihalo-alkanes, for instance bromo chloroethane, and the reaction is

5 preferably carried out in an organic solvent such as, for example, dimethylformamide, with a strong base such as sodium hydride, at a temperature of from 0° C. to 100° C., for instance at room temperature.

10 Compounds of formula (VII) above can be prepared, for example, as follows:

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As mentioned above, the compounds of the invention and
5 their pharmaceutically acceptable salts have useful
central nervous system activity. The compounds are
active at the serotonin, 5-HT2A, receptor. Their
binding activity has been demonstrated in a test
described by Nelson, D. L. et al, J. Pharmacol. Exp.
10 Ther., 265, 1272-1279, in which the affinity of the
compound for the human 2A receptor is measured by its
ability to displace the ligand [³H] ketanserine. In
this test, the compounds of the invention in the
following Examples had a Ki of less than 15 nM. The
15 compounds of the invention are also active serotonin

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reuptake inhibitors as measured by their displacement of [³H] paroxetine at the reuptake site, *Neuropharmacology* Vol. 32 No. 8, 1993, pages 737-743.

5 Because of their selective affinity for 5-HT receptors, the compounds of the present invention are indicated for use in treating a variety of conditions such as depression, obesity, bulimia, alcoholism, pain, hypertension, ageing, memory loss, sexual dysfunction, 10 anxiety, schizophrenia, gastrointestinal disorders, headache, cardiovascular disorders, smoking cessation, drug addiction, emesis, Alzheimer's and sleep disorders.

The compounds of the invention are effective over a wide 15 dosage range, the actual dose administered being dependent on such factors as the particular compound being used, the condition being treated and the type and size of mammal being treated. However, the dosage required will normally fall within the range of 0.01 to 20 20 mg/kg per day, for example in the treatment of adult humans, dosages of from 0.5 to 100 mg per day may be used.

The compounds of the invention will normally be 25 administered orally or by injection and, for this purpose, the compounds will usually be utilised in the

form of a pharmaceutical composition. Such compositions are prepared in a manner well known in the pharmaceutical art and comprise at least one active compound.

5

Accordingly the invention includes a pharmaceutical composition comprising as active ingredient a compound of formula (I) or a pharmaceutically acceptable salt or ester thereof, associated with a pharmaceutically acceptable excipient. In making the compositions of the invention, the active ingredient will usually be mixed with a carrier, or diluted by a carrier, or enclosed within a carrier which may be in the form of a capsule, sachet, paper or other container. The excipient may be a solid, semi-solid or liquid material which acts as a vehicle, excipient or medium for the active ingredient. Some examples of suitable excipients are lactose, dextrose, sucrose, sorbitol, mannitol, starches, gum acacia, calcium phosphate, alginates, tragacanth, gelatin syrup, methyl cellulose, methyl- and propyl-hydroxybenzoate, talc, magnesium stearate or oil. The compositions of the invention may, if desired, be formulated so as to provide quick, sustained or delayed release of the active ingredient after administration to the patient.

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Depending on the route of administration, the foregoing compositions may be formulated as tablets, capsules or suspensions for oral use and injection solutions or suspensions for parenteral use or as suppositories.

5 Preferably the compositions are formulated in a dosage unit form, each dosage containing from 0.5 to 100 mg, more usually 1 to 100 mg, of the active ingredient.

10 The following Preparations and Examples illustrate routes to the synthesis of the compounds of the invention.

PREPARATION 1

15 6-Fluoroindole

1-Dimethylamino-2-(4-fluoro-2-nitro)phenylethene

A mixture of 4-fluoro-2-nitrotoluene (50 g, 0.32 mol), dimethylformamide dimethylacetal (76.77 g) and dimethylformamide (910 ml) were 20 heated under reflux under nitrogen with stirring for 7 hours, cooled, allowed to stand for 16 hours, poured into ice-water (2000 ml), stirred for 15 minutes and the resultant precipitate isolated by filtration, washed with water (500 ml), dried to 25 give a red solid.

6-Fluoroindole

A 40 litre Cook hydrogenator was charged under a nitrogen atmosphere with 10% palladium on charcoal (9 g) suspended in toluene (400 ml). To this suspension was added 1-dimethylamino-2-(4-fluoro-2-nitro)phenylethene (137.2 g, 0.653 mol) in toluene (1400 ml) and the mixture hydrogenated at 80 psi for 3.5 hours. The suspension was then filtered through a celite pad, which was washed through with toluene (2 x 200 ml) and the filtrate and washings evaporated under reduced pressure to give a brown oil which crystallised on standing to a yellow brown solid 93.65 g. This solid was dissolved in ethyl acetate-hexane (7:3) and filtered through a pad of flash silica. The required fractions were collected and evaporated under reduced pressure to give a pale brown solid.

20 PREPARATION 24-(6-Fluoroindol-3-yl)-1,2,5,6-tetrahydropyridine

Powdered potassium hydroxide (144.4 g) was added carefully to a mechanically stirred mixture of 6-fluoroindole (49.23 g, 0.364 mol) and

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4-piperidone monohydrate (111.93 g, 0.728 mol) in methanol (1500 ml). The mixture was then heated under reflux under nitrogen for 18 hours and then more potassium hydroxide (40 g) was added and the 5 reaction mixture heated under reflux for a further 4 hours. The reaction mixture was allowed to cool to room temperature and poured onto ice-water (3000 ml) and stirred for 1 hour and the precipitated solid isolated by filtration and dried 10 at 50° C. in vacuo to give a solid.

EXAMPLE 1

15 1-[2-[4-(6-fluoroindol-3-yl)-1,2,5,6-tetrahydro-1-pyridyl]-1-ethyl]-3-methyl-1,3-dihydro-2,1,3-benzothiadiazole-2,2-dioxide

To a 250 ml 3-necked round bottom flask equipped with 20 reflux condenser, thermometer, magnetic stirrer bar and nitrogen bubbler was charged with sulphonamide (9.61 g; 0.1 mol) and pyridine (100 ml) and the stirred solution heated to reflux under nitrogen. N-methyl-1,2-phenylene diamine (12.2 g, 0.1 mol) in dry pyridine (30 ml) was 25 added dropwise to the solution whilst maintaining reflux. After 5 hours, the reaction mixture allowed to

cool and the pyridine removed under reduced pressure. The residue was dissolved in 5N hydrochloric acid (100 ml) and ethyl acetate (100 ml) and the acidic layer was extracted with further ethyl acetate (5 x 100 ml).

5 The combined organic layer was washed with 5N hydrochloric acid (2 x 100 ml), extracted with 2N sodium hydroxide solution (3 x 100 ml) and the combined aqueous layer washed with diethyl ether (2 x 150 ml). Ice was then added followed by the addition of 5N hydrochloric

10 acid with cooling and stirring of the suspension to pH 1. The oily suspension was stirred for several hours at room temperature when a colourless solid separated. The solid was filtered and dried at room temperature under vacuum to leave a light pink solid, 1,3-dihydro-1-methyl-2,1,3-benzothiadiazole-2,2-dioxide, which was

15 used directly in the next step.

1,3-dihydro-1-methyl-2,1,3-benzothiadiazole-2,2-dioxide (1.36 g, 7.4 mmol) was dissolved in DMF (40 ml) and then

20 treated with sodium hydride (0.33 g, 60% oil dispersion, 8.2 mmol, 1.1 equivalent). The mixture was stirred at room temperature and under nitrogen for 45 minutes. 1-Bromo-2-chloroethane (0.74 ml, 1.27 g, 8.9 mmol, 1.2 equivalent) was added in one portion to the stirred

25 mixture, and stirred overnight at room temperature. The solvent was removed *in vacuo* and the residue suspended

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in water and extracted into ethyl acetate (3 x 40 ml). The bulk extracts were washed with water (3 x 50 ml) and brine, then dried over anhydrous magnesium sulfate. Filtration was followed by evaporation to dryness 5 *in vacuo* and the residue chromatographed on silica using dichloromethane as eluent. This gave a white solid [1-(2-chloroethyl)-1,3-dihydro-3-methyl-1H-2,1,3-benzothiadiazole-2,2-dioxide].

10 A mixture of 4-(6-fluoroindol-3-yl)-1,2,5,6-tetrahydropyridine (0.87 g, 4.0 mmol, 1.05 equivalent), 1-(2-chloroethyl)-1,3-dihydro-3-methyl-1H-2,1,3-benzothiadiazole-2,2-dioxide (1.1 g, 4.46 mmol), anhydrous sodium carbonate (2.34 g, 22.3 mmol, 5 equivalents), sodium iodide (0.67 g; 4.46 mmol) and de-ionised water (20 ml) was rapidly stirred and warmed under reflux for 20 hours. After cooling to room temperature, the mixture was extracted with chloroform (3 x 30 ml). The bulked extracts were washed with water 15 and then dried over magnesium sulfate. Filtration was followed by evaporation to dryness *in vacuo* to yield an orange solid. This material was purified further by chromatography on silica using dichloromethane initially followed by ethyl acetate to give the final product as 20 an orange solid which was triturated with a mixture of diethyl ether and ethyl acetate. This gave a yellow 25

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solid after filtration, 1-{2-[4-(6-fluoroindol-3-yl)-
1,2,5,6-tetrahydro-1-pyridyl]-1-ethyl}-3-methyl-1,3-
dihydro-2,1,3-benzothiadiazole-2,2-dioxide.

5 The compound was converted into its hydrochloride salt
using ethereal HCl in ethanol with M.P. 246-8° C.

10 The following Examples illustrate typical formulations
containing the compound of the invention.

EXAMPLE 2

Tablets each containing 10 mg of active ingredient are
15 made up as follows:

Active ingredient	10 mg
Starch	160 mg
Microcrystalline cellulose	100 mg
Polyvinylpyrrolidone (as 10% solution in water)	13 mg
20 Sodium carboxymethyl starch	14 mg
Magnesium stearate	3 mg
<hr/>	
Total	300 mg
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- 20 -

The active ingredient, starch and cellulose are mixed thoroughly. The solution of polyvinylpyrrolidone is mixed with the resultant powders and passed through a sieve. The granules so produced are dried and re-passed 5 through a sieve. The sodium carboxymethyl starch and magnesium stearate are then added to the granules which, after mixing, are compressed on a tablet machine to yield tablets each weighing 300 mg.

10 EXAMPLE 3

Capsules each containing 20 mg of medicament are made as follows:

15	Active ingredient	20 mg
	Dried starch	178 mg
	Magnesium stearate	2 mg

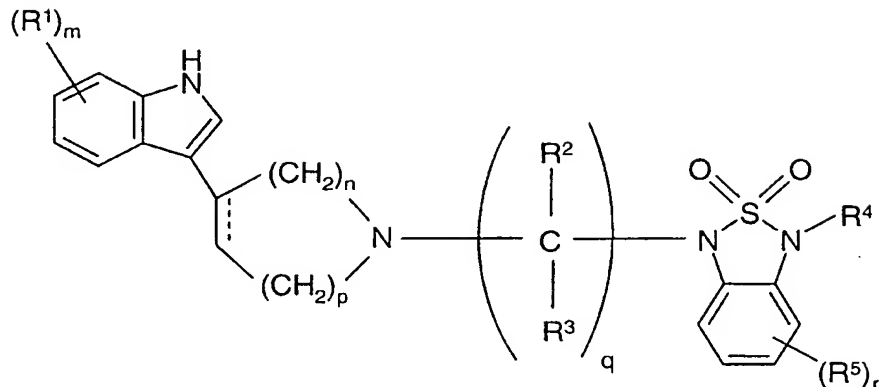
20	Total	200 mg

The active ingredient, starch and magnesium stearate are passed through a sieve and filled into hard gelatine capsules in 200 mg quantities.

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CLAIMS

1. A compound of the following formula:



5

in which m is 0, 1 or 2, n is 1 or 2, p is 1 or 2,

q is 1 to 6, r is 0 or 1 to 3,

10

R^1 is halo, C_{1-4} alkyl, nitrile, trifluoromethyl or
 C_{1-4} alkoxy,

R^2 and R^3 are each hydrogen or C_{1-4} alkyl,

15

R^4 is hydrogen, C_{1-4} alkyl, optionally substituted phenyl or optionally substituted phenyl- C_{1-4} alkyl,

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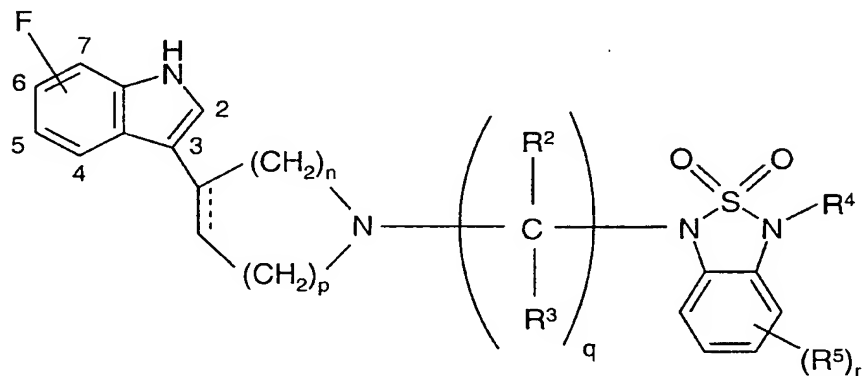
R^5 is C_{1-4} alkyl, C_{1-4} alkoxy, carboxy, hydroxy, cyano, halo, trifluoromethyl, nitro or amino, and

the dotted line represents an optional double bond;

5

and salts and esters thereof.

2. A compound according to Claim 1 of the formula:



in which n is 1 or 2, p is 1 or 2,

q is 1 to 6, r is 0 or 1 to 3,

15

R^2 and R^3 are each hydrogen or C_{1-4} alkyl,

R^4 is hydrogen, C_{1-4} alkyl, optionally substituted phenyl or optionally substituted phenyl- C_{1-4} alkyl,

R⁵ is C₁₋₄ alkyl, C₁₋₄ alkoxy, carboxy, hydroxy, cyano, halo, trifluoromethyl, nitro or amino,

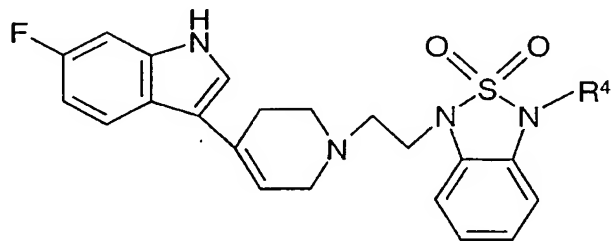
5 the dotted line represents an optional bond, and

the fluorine atom is attached at the 6 or
7-position;

10 or a salt or ester thereof.

3. A compound according to Claim 2, in which the
dotted line represents a double bond, n is 2 and p
is 1, R² and R³ are both hydrogen, q is 2, R⁴ is
15 C₁₋₄ alkyl, and r is 0 or 1.

4. A compound according to Claim 2 of the formula:



20

in which R⁴ is C₁₋₄ alkyl.

5. A pharmaceutical formulation comprising a compound as defined in any of Claims 2 to 4, or a pharmaceutically acceptable salt or ester thereof, together with a pharmaceutically acceptable diluent or carrier therefor.
10. A compound according to Claim 1, or a pharmaceutically acceptable salt or ester thereof, for use as a pharmaceutical.
15. Use of a compound according to any of Claims 1 to 4, or a pharmaceutically acceptable salt or ester thereof, in the manufacture of a medicament for use in the treatment of a disorder of the central nervous system.

INTERNATIONAL SEARCH REPORT

Int. Application No
PCT/GB 00/00469

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07D417/14 A61K31/41 A61P25/00		
According to International Patent Classification (IPC) or to both national classification and IPC		
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07D A61K		
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched		
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)		
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	EP 0 013 612 A (JANSSEN PHARMACEUTICA N.V.) 23 July 1980 (1980-07-23) claims 1-13 ---	1-7
Y	EP 0 184 258 A (JANSSEN PHARMACEUTICA N.V.) 11 June 1986 (1986-06-11) claims 1-10 ---	1-10
P, Y	EP 0 897 921 A (ELI LILLY AND COMPANY LIMITED) 24 February 1999 (1999-02-24) claims 1-12 ---	1-10
Y	EP 0 058 975 A (BOEHRINGER INGELHEIM LTD.) 1 September 1982 (1982-09-01) claims 1-7 ---	1-10 -/-
<input checked="" type="checkbox"/> Further documents are listed in the continuation of box C.		<input checked="" type="checkbox"/> Patent family members are listed in annex.
* Special categories of cited documents : "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed		
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Date of the actual completion of the international search		Date of mailing of the international search report
28 April 2000		09/05/2000
Name and mailing address of the ISA European Patent Office, P.B. 5518 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016		Authorized officer Herz, C

INTERNATIONAL SEARCH REPORT

International Application No
PCT/GB 00/00469

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
P, Y	WO 99 14203 A (TAKEDA CHEMICAL INDUSTRIES, LTD.) 25 March 1999 (1999-03-25) claims 1-26 ----	1-10
Y	EP 0 433 149 A (RHONE-POULENC SANTE) 19 June 1991 (1991-06-19) cited in the application claims 1- ----	1-10
Y	EP 0 854 146 A (ELI LILLY AND COMPANY LTD.) 22 July 1998 (1998-07-22) claims 1- ----	1-10
Y	FR 2 621 588 A (ROUSSEL-UCLAF) 14 April 1989 (1989-04-14) claims 1-12 -----	1-10

INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No
PCT/GB 00/00469

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
EP 13612	A 23-07-1980	US 4335127 A		15-06-1982
		AT 5258 T		15-11-1983
		AU 536175 B		19-04-1984
		AU 5438180 A		17-07-1980
		CA 1132557 A		28-09-1982
		CS 223977 B		25-11-1983
		CY 1252 A		31-08-1984
		DE 3065489 D		15-12-1983
		DK 7280 A		09-07-1980
		ES 487537 D		16-12-1980
		ES 8101586 A		16-03-1981
		FI 800047 A, B,		09-07-1980
		GR 67304 A		29-06-1981
		HK 76284 A		19-10-1984
		HR 930474 B		31-08-1997
		HU 184222 B		30-07-1984
		IE 49351 B		18-09-1985
		IL 59084 A		29-02-1984
		JP 1497429 C		29-05-1989
		JP 55105679 A		13-08-1980
		JP 63046753 B		19-09-1988
		KR 8500684 B		14-05-1985
		KR 8500677 B		10-05-1985
		KR 8500683 B		14-05-1985
		LU 88219 A		03-02-1994
		MY 55587 A		31-12-1987
		NO 800034 A, B,		09-07-1980
		NZ 192551 A		06-07-1984
		PH 17114 A		01-06-1984
		PL 221249 A		01-12-1980
		PT 70662 A		01-02-1980
		RO 79148 A		17-08-1982
		SG 48684 G		29-03-1985
		SI 8010046 A		31-12-1994
		SU 1041034 A		07-09-1983
		YU 4680 A		30-06-1983
		ZA 8000082 A		26-08-1981
		US 4522945 A		11-06-1985
EP 184258	A 11-06-1986	AT 75741 T		15-05-1992
		CA 1246074 A		06-12-1988
		DE 3585990 A		11-06-1992
		JP 2009087 C		11-01-1996
		JP 7045492 B		17-05-1995
		JP 61148175 A		05-07-1986
		US 4665075 A		12-05-1987
EP 897921	A 24-02-1999	US 5929072 A		27-07-1999
		AU 8743098 A		16-03-1999
		CA 2245331 A		22-02-1999
		WO 9910346 A		04-03-1999
		JP 11116572 A		27-04-1999
EP 58975	A 01-09-1982	US 4359468 A		16-11-1982
		AR 228475 A		15-03-1983
		AT 10742 T		15-12-1984
		AU 543948 B		09-05-1985
		AU 8078382 A		02-09-1982

INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No
PCT/GB 00/00469

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
EP 58975	A	CA 1191137 A		30-07-1985
		CS 227343 B		16-04-1984
		DD 202562 A		21-09-1983
		DE 3261497 D		24-01-1985
		DK 79882 A, B,		26-08-1982
		ES 509871 D		01-05-1983
		ES 8306142 A		01-08-1983
		ES 517988 D		01-01-1984
		ES 8401961 A		01-04-1984
		FI 820594 A, B,		26-08-1982
		GB 2093455 A, B		02-09-1982
		GR 74778 A		12-07-1984
		HU 187652 B		28-02-1986
		IE 52562 B		09-12-1987
		JP 1654161 C		13-04-1992
		JP 3018637 B		13-03-1991
		JP 57156484 A		27-09-1982
		KR 8900008 B		02-03-1989
		NO 820583 A, B,		26-08-1982
		NZ 199822 A		31-01-1985
		PH 17889 A		21-01-1985
		PL 235187 A		26-03-1984
		PT 74481 A, B		01-03-1982
		SU 1088665 A		23-04-1984
		YU 41082 A		20-03-1985
		ZA 8201196 A		26-10-1983
WO 9914203	A	25-03-1999	AU 9002598 A	05-04-1999
			JP 11152275 A	08-06-1999
EP 433149	A	19-06-1991	FR 2655652 A	14-06-1991
			FR 2662696 A	06-12-1991
			AT 101612 T	15-03-1994
			AU 643241 B	11-11-1993
			AU 6798190 A	20-06-1991
			CA 2032104 A	14-06-1991
			DE 69006699 D	24-03-1994
			DE 69006699 T	09-06-1994
			DK 433149 T	14-03-1994
			ES 2062465 T	16-12-1994
			FI 906108 A	14-06-1991
			HU 56566 A, B	30-09-1991
			JP 3255063 A	13-11-1991
			NO 905368 A	14-06-1991
			NZ 236436 A	26-03-1993
			PT 96197 A	30-09-1991
			US 5130313 A	14-07-1992
			ZA 9009982 A	30-10-1991
EP 854146	A	22-07-1998	AT 190617 T	15-04-2000
			AU 5569298 A	07-08-1998
			CA 2227132 A	17-07-1998
			CZ 9901510 A	15-09-1999
			DE 69800092 D	20-04-2000
			WO 9831686 A	23-07-1998
			JP 10204090 A	04-08-1998
			NO 993372 A	08-07-1999
			ZA 9800300 A	10-09-1998

INTERNATIONAL SEARCH REPORT

Information on patent family members

Int. Application No
PCT/GB 00/00469

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
FR 2621588	A 14-04-1989	NONE	